PATENT COOPERATION TREATY

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INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

(Chapter II of the Patent Cooperation Treaty)

(PCT Article 36 and Rule 70)

Applicant's or agent's P36154A/EBA/B0		FOR FURTHER A	CTION	See Form PCT/IPEA/416
International applicat PCT/GB2005/000		International filing date 14.02.2005	day/month/year)	Priority date (day/month/year) 12.02.2004
International Patent (INV. C12N5/06	Classification (IPC) or n	ational classification and	IPC	
Applicant UNIVERSITY OF	NEWCASTLE UF	ON TYNE et al.		
		liminary examination r nsmitted to the applica		this International Preliminary Examining e 36.
2. This REPOR	T consists of a total of	of 8 sheets, including	this cover sheet.	
3. This report is	also accompanied b	y ANNEXES, compris	ing:	
a.⊠ sent to	o the applicant and t	o the International Bur	eau) a total of 3 shee	ets, as follows:
aı		ng rectifications authoi		n amended and are the basis of this report (see Rule 70.16 and Section 607 of the
be	neets which supersections which supersections which supplemental box.	de earlier sheets, but v in the international ap	which this Authority co plication as filed, as in	onsiders contain an amendment that goes ndicated in item 4 of Box No. I and the
seque	nce listing and/or tab		celectronic form only,	nber of electronic carrier(s)), containing a as indicated in the Supplemental Box structions).
4. This report co	ntains indications re	lating to the following i	tems:	
⊠ Box No. I	Basis of the rep	ort		
☐ Box No. II	Priority			
⊠ Box No. II	l Non-establishm	ent of opinion with reg	ard to novelty, inventi	ve step and industrial applicability
⊠ Box No. I'	/ Lack of unity of	invention		
⊠ Box No. V		ment under Article 35(ations and explanations	, —	elty, inventive step or industrial tement
☐ Box No. V	l Certain docume	nts cited		
□ Box No. V	Il Certain defects	in the international app	lication	
☐ Box No. V	III Certain observa	tions on the internatior	nal application	
Date of submission of	the demand		Date of completion of	f this report
26.01.2006			28.03.2006	
	dress of the internation	al	Authorized officer	hes Palenta.
preliminary examining authority: European Patent Office D-80298 Munich Tel. +49 89 2399 - 0 Tx: 523656 epmu d Fax: +49 89 2399 - 4465			Nichogiannopoule Telephone No. +49 8	ou, A

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/GB2005/000518

	Box No. I Basis of the report	
1.	With regard to the language, thi filed, unless otherwise indicated	is report is based on the international application in the language in which it was under this item.
	which is the language of a t international search (und publication of the interna	slations from the original language into the following language, ranslation furnished for the purposes of: der Rules 12.3 and 23.1(b)) ational application (under Rule 12.4) examination (under Rules 55.2 and/or 55.3)
2.		the international application, this report is based on (replacement sheets which iving Office in response to an invitation under Article 14 are referred to in this e not annexed to this report):
	Description, Pages	
	1-37	as originally filed
	Sequence listings part of the des	cription, Pages
	1-3	as originally filed
	Claims, Numbers	
	1-13	received on 30.01.2006 with letter of 26.01.2006
	Drawings, Sheets	
	1/8-8/8	as originally filed
	a sequence listing and/or an	y related table(s) - see Supplemental Box Relating to Sequence Listing
3.	☐ The amendments have result the description, pages ☐ the claims, Nos. ☐ the drawings, sheets/figs ☐ the sequence listing (specially any table(s) related to se	ecify):
1.	•	ecify):
	* If item 4 applies, so	me or all of these sheets may be marked "superseded."

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/GB2005/000518

	Box No. III Non-establishment of opinion with regard to novelty, inventive step and industrial applicability				
	he questions whether the claimed invention appears to be novel, to involve an inventive step (to be non- bvious), or to be industrially applicable have not been examined in respect of:				
	the entire international applica	tion,			
\boxtimes	claims Nos. 5-13				
	because:				
	the said international application not require an international pre		the said claims Nos. relate to the following subject matter which does ary examination (specify):		
	the description, claims or draw that no meaningful opinion cou		(indicate particular elements below) or said claims Nos. are so unclear formed (specify):		
	the claims, or said claims Nos. are so inadequately supported by the description that no meaningful opinion could be formed.				
\boxtimes	no international search report h	nas b	een established for the said claims Nos. 5-13		
	the nucleotide and/or amino acid sequence listing does not comply with the standard provided for in Annex C of the Administrative Instructions in that:				
	the written form		has not been furnished		
			does not comply with the standard		
	the computer readable form		has not been furnished		
			does not comply with the standard		
	the tables related to the nucleo not comply with the technical re	tide a equir	and/or amino acid sequence listing, if in computer readable form only, do ements provided for in Annex C-bis of the Administrative Instructions.		
	See separate sheet for further	detai	ls		

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/GB2005/000518

	Box	No. IV	Lack of unity of inve	ention				
1.		☐ restri ☐ paid ☐ paid	onse to the invitation to icted the claims. additional fees. additional fees under per restricted nor paid a	orotest	•	itional fees, the app	olicant has:	
2.		This Aut Rule 68	thority found that the re .1, not to invite the app	equirer dicant	nent of unity to restrict or	of invention is not pay additional fees	complied with and chose, according s.	j to
3.	This	s Authorit	ty considers that the re	quiren	nent of unity	of invention in acco	ordance with Rules 13.1, 13.2 and 1	3.3
		complie	d with.					
	\boxtimes	not com	plied with for the follov	ving re	asons:			
		see sep	parate sheet					
4.	Cor	nsequent	ly, this report has beer	estab	lished in res	pect of the followin	g parts of the international application	on:
		all parts	i.			•		
	\boxtimes	the part	s relating to claims No	s. 1-4				
		« No. V olicability	Reasoned statemer y; citations and expla	nt und ination	er Article 35 ns supporti	5(2) with regard to ng such statement	novelty, inventive step or industr	rial
1.	Sta	tement						
	Nov	elty (N)		Yes: No:	Claims Claims	1-4		
	Inve	entive ste	ep (IS)	Yes: No:	Claims Claims	1-4		
	Ind	ustrial ap	plicability (IA)	Yes: No:	Claims Claims	1-4		
2.	Cita	ations and	d explanations (Rule 7	0.7):				
	see	e separat	te sheet				e some en	

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY

International application No. PCT/GB2005/000518

Supplemental Box relating to Sequence Listing

C	711 (11	iua	tion of box i, item 2.		
1.	With regard to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the claimed invention, this report has been established on the basis of:				
	a. ty	ype	of material:		
		\boxtimes	a sequence listing		
			table(s) related to the sequence listing		
	b. fo	orm	at of material:		
		\boxtimes	in written format		
		\boxtimes	in computer readable form		
	c. time of filing/furnishing:				
	٥	\boxtimes	contained in the international application as filed		
		\boxtimes	filed together with the international application in computer readable form		
			furnished subsequently to this Authority for the purposes of search and/or examination		
			received by this Authority as an amendment on		
2.		the add	addition, in the case that more than one version or copy of a sequence listing and/or table(s) relating creto has been filed or furnished, the required statements that the information in the subsequent or ditional copies is identical to that in the application as filed, appropriate, were furnished.		

3. Additional observations, if necessary:

Re Item I

Basis of the report

1. The amendments filed with the letter of 26.01.2006 are formally allowable under Article 34(2)(b) PCT because they do not introduce subject-matter extending beyond the content of the application as filed.

Re Item II

Priority

1. The present application validly claims priority from 12.02.2004. Any documents cited in the International Search Report as P documents have therefore not been considered as comprised in the prior art relevant for the present application.

Re Item III

Non-establishment of opinion with regard to novelty, inventive step and industrial applicability

1. No meaningful examination could be performed for new claims 5-13, for the following reason:

No complete international search report has been established for said claims, corresponding to original claims 10-14, 22 and 17-19 (see Form PCT/ISA/210). Accordingly, said claims need not be the subject of international preliminary examination (Rule 66. 1.(e) (PCT)).

Re Item IV

Lack of unity of invention

1. The IPEA agrees with the objection put forward by the ISA as to lack of unity pursuant to Rule 13 PCT, and considers that the present invention (new claims 1-13)

relates to three distinct groups of inventions. New claims 1-13 correspond to original claims 6-14, 22 and 17-19, which belong to three distinct groups of inventions (groups I, II and III) for the reasons outlined in Form PCT/ISA/210.

Re Item V

Reasoned statement with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

- 1. Reference is made to the following documents:
 - D1: RICHARDS M et al.: "Human feeders support prolonged undifferentiated growth of human inner cell masses and embryonic stem cells" NATURE BIOTECHNOLOGY, vol. 20, no. 9, September 2002, pages 933-936
 - D2: HOVATTA O et al.: "A culture system using human foreskin fibroblasts as feeder cells allows production of human embryonic stem cells." HUMAN REPRODUCTION, vol. 18, no. 7, July 2003 (2003-07), pages 1404-1409,
 - D3: HENDERSON J K et al.: "Preimplantation human embryos and embryonic stem cells show comparable expression of stage-specific embryonic antigens." STEM CELLS 2002, vol. 20, no. 4, 2002, pages 329-337,
- 2. Novelty and Inventive step (Article 33(2) and (3) PCT)
- 2.1. The present application (Invention I, new claims 1-4) discloses the human embryonic stem cell line hES-NCL1, a stem cell bank comprising it and methods for screening agents for toxicity using it.
- 2.2. **D1** is a publication disclosing the derivation of a new human ES cell line with the Oct-4, SSEA-4, Tra1-60 and GCTM-2 phenotype.
 - **D2** is a publication disclosing the culture of huES cells on human foreskin fibroblasts, having the Oct-4, SSEA-4, Tra1-60 phenotype.
 - **D3** is a publication disclosing that huES cells express SSEA3, SSEA4, TRA-1-60, Oct-4 and Rex1.
- 2.3. None of the available prior art discloses the specific deposited cell-line of new claim 1. Said claim as well as claims 2-4 referring to it are thus considered novel and

INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (SEPARATE SHEET)

International application No.

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inventive under the terms of Articles 33(2) and (3) PCT.

3. Industrial applicability (Article 33(4) PCT)

The subject-matter of the claims for which an opinion has been established (see item III) appears to be industrially applicable under the terms of Article 33(4) PCT.

Re Item VIII

Certain observations on the international application

1. Applicant's attention is drawn to the fact that, upon entry into the regional phase, patentability of claims relating to human embryos may underlie restrictions based on moral grounds. The EPO, for example, does not recognize as patentable the subject-matter of claims to the cloning of human beings, the modification of the germ line identity of human beings and the use of human embryos for industrial or commercial purposes (Article 53(a) and Rule 23d EPC). Claims to human embryonic stem cells might be regarded as falling under said exclusions.

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1	Claims	·
2		
3	1. The	e stem cell line hES-NCL1 deposited at
4	NIE	SSC under Accession No. P-05-001.
5		
6	2. An	embryonic stem cell bank comprising a
7	mul	tiplicity of genetically distinct stem
8	cel	l lines, including the stem cell line as
9	cla	imed in Claim 1.
10		
11	3. A m	ethod of screening an agent for toxicity
12	and	or for therapeutic efficacy, said method
13	com	prising:
14	i.	exposing the stem cell line as claimed in
15		Claim 1 to said agent;
16	11.	monitoring any alteration in viability
17		and/or metabolism of said stem cells; and
18	iii.	determining any toxic or therapeutic
19		effect of said agent.
20		
21	4. A me	ethod of screening an agent for toxicity
22	and	or for therapeutic efficacy, said method
23	comp	prising:
24	i.	exposing an embryonic stem cell bank as
25		claimed in Claim 2 to said agent;
26	ii.	monitoring any alteration in viability
27		and/or metabolism of said stem cells;
28		and
29	iii.	determining any toxic or therapeutic
30		effect of said agent.
31		

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		39
1	5.	A method of producing fibroblast-like cells,
2		said method comprising:
3		i. providing the stem cell line as claimed
4		in Claim 1;
5		ii. allowing cells of said stem cell line to
6		differentiate into stem cell derived
7		fibroblast-like cells.
8		
9	6.	The method of Claim 5 which is conducted
10		without use of a specific stimulant for
11		differentiation.
12		e
13	7.	The method as claimed in either one of Claims
14		5 and 6 wherein the fibroblast-like cells are
15		produced for a therapeutic purpose.
16		
17	8.	A method of culturing cells wherein the
18		fibroblast-like cells obtained as claimed in
19		Claims 5 or 6 act as feeder cells or
20		condition cell culture media used during
21		culture of the cells.
22		
23	9.	The method as claimed in Claim 8 wherein the
24		cells being cultured are stem cells.
25		
26	10.	A self-feeder system for the growth of
27		undifferentiated stem cells, said system
28		comprising:
29		i. culturing the stem cell line as claimed
30		in Claim 1; and
31		ii. allowing some of the cells of said stem
32		cell line to differentiate into stem

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1		cell derived fibroblast-like cells
2		whilst the remainder of the cells of
3		said embryonic stem cell line remain in
4		an undifferentiated pluripotent,
5		multipotent or unipotent state, whereby
6		said stem cell derived fibroblast-like
7		cells act as autogeneic feeder cells for
8		said stem cells.
9		
10	11.	The fibroblast-like cell line hESCdF-NCL as
11		deposited at ECACC under Accession No.
12		04010601.
13		
14	12.	A method of culturing cells wherein hESCdF-
15		NCL cells act as feeder cells or condition
16		cell culture media used during culture of the
17		cells.
18		
19	13.	The method as claimed in Claim 12 wherein the
20		cells being cultured are stem cells.